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- (56) Documents Cited WO 95/16007 A1
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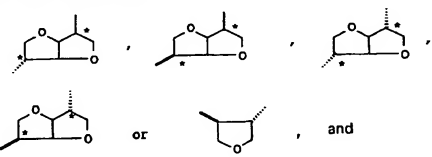
(54) Abstract Title

Polymerizable chiral dopants based on chiral tetrahydrofuran or hexahydrofuro[3,2-b]furan substituted by at least two mesogenic groups

Chiral compounds are disclosed of the formula

where, independently of one another,

- Α
- is a mesogenic group containing two phenylene radicals which are unsubstituted or substituted by C₁-M to C4-alkyl, methoxy, ethoxy, fluorine, chlorine, bromine, C1- to C20-alkoxycarbonyl or C1- to C₂₀-alkylcarbonyl and are linked via O, CO, OCO, OCOO or COO,
- is a direct bond, O, S, COO, OCO, OCOO, CON(R) or N(R)CO, and
- is a polymerizable group, Z
- m is 0 or 1,
- is from 2 to 6, and
- is a chiral radical of the formula



is C1- to C4-alkyl or hydrogen,

and their use as polymerizable, chiral high-twisting-power dopants for the production of cholesteric networks. These compounds are suitable for use in electro-optical displays or as chiral dopants for nematic or cholesteric liquid crystals for producing layers which reflect in color. They may also be used as chiral dopants in optical polarizers and optical filters.

Polymerizable chiral compounds, and their use

- It is known that molecules which are anisotropic in shape can form liquid-crystalline phases, known as mesophases, on warming. The individual phases differ through the spatial arrangement of the major parts of the molecules on the one hand and through the molecular arrangement with respect to the long axes on the other hand (G.W. Gray, P.A. Winsor, Liquid Crystals and Plastic
- 10 Crystals, Ellis Horwood Limited, Chichester 1974). The nematic liquid-crystalline phase is distinguished by the fact that there is only one alignment long-distance ordering due to the long molecular axes lining up in parallel. Under the prerequisite that the molecules making up the nematic phase are chiral, a
- 15 cholesteric phase forms, in which the long axes of the molecules form a helical superstructure perpendicular thereto (B. Baessler, Festkörperprobleme XI, 1971). The chiral moiety may be present either in the liquid-crystalline molecule itself or added to the nematic phase as a dopant. Phases produced by doping are referred
- 20 to as induced cholesteric phases. This phenomenon was first studied on cholesterol derivatives (H. Baessler, M.M. Labes, J. Chem. Phys. 52 (1970) 631; H. Beassler, T.M. Laronge, M.M. Labes. J. Chem. Phys. 51 (1969) 3213; H. Filkelmann, H. Stegemeyer, Z. Naturforschg. 28a (1973) 799). It later also became possible to
- 25 induce cholesteric phases by adding other chiral substances which are not themselves liquid-crystalline (H. Stegemeyer, K.J. Mainusch, Naturwiss. 58 (1971) 599; H. Finkelmann, H. Stegemeyer, Ber. Bunsenges. Phys. Chem. 78 (1974) 869).
- 30 The cholesteric phase has remarkable optical properties: large optical rotation and pronounced circular dichroism caused by selective reflection of circular-polarized light within the cholesteric layer. The different colors observed depending on the veering angle depend on the pitch of the helical superstructure,
- 35 which is itself dependent on the twisting power of the chiral component. The pitch and thus the wavelength range of the selectively reflected light of a cholesteric layer can be varied, in particular, by changing the concentration of a chiral dopant (J.E. Adams, W.E.L. Haas, Mol. Cryst. Liq. Cryst. 16 (1972) 33).
- 40 Such cholesteric systems offer interesting opportunities for practical use. Thus, incorporation of chiral moieties into mesogenic acrylic esters after alignment in the cholesteric phase and photocrosslinking allows the production of a stable, colored network, but the concentration of the chiral component therein
- 45 cannot be changed (G. Galli, M. Laus, A. Angeloni, Makromol. Chem. 187 (1986) 289). Furthermore, admixing of non-crosslinkable, chiral compounds with nematic acrylic esters

allows, after photocrosslinking, the production of a colored polymer (I. Heyndericks, D.J. Broer, Mol. Cryst. Liq. Cryst. 203 (1991) 113), but this still contains volatile constituents which are prohibitive for use.

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EP-A-739403 describes chiral compounds which firstly have a high twisting power and secondly can be incorporated into the cholesteric phase in a stable manner in any desired concentration without diffusing or crystallizing out of the phase.

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We have now found a group of compounds which has even better properties than those described above.

Accordingly, the present invention provides compounds of the 15 formula I

$$(Z-Y-[A]_m-Y-M-Y-)_n X I,$$

where, independently of one another,

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- A is a spacer,
- M is a mesogenic group containing two phenylene radicals which are unsubstituted or substituted by C₁- to C₄-alkyl, methoxy, ethoxy, fluorine, chlorine, bromine, C₁- to C₂₀-alkoxycarbonyl or C₁- to C₂₀-alkylcarbonyl and are linked via O, CO, OCO, OCOO or COO,
- Y is a direct bond, O, S, COO, OCO, OCOO, CON(R) or N(R)CO, and
- z is a polymerizable group,
- 30 m is 0 or 1,
 - n is from 2 to 6, and
 - X is a chiral radical of the formula

45

R is C_1 - to C_4 -alkyl or hydrogen.

The spacer A can be any group known for this purpose; the spacers are usually linked to X via ester or ether groups or a direct bond. The spacers generally contain from 1 to 30, preferably from 1 to 12, carbon atoms and may be interrupted in the chain by, for example, O, S, NH or NCH₃. Suitable substituents for the spacer chain are fluorine, chlorine, bromine, cyano, methyl and ethyl.

Examples of representative spacers are the following:

10 (CH₂)_p, (CH₂CH₂O)_qCH₂CH₂,

$$CH_3$$
 CH_3 CH_3 CH_3 CH_3 CH_2 CH_2 CH_2 CH_2 CH_3 CH_3

15

where

p is from 1 to 12, and

q is from 1 to 3.

20 Particularly preferred mesogenic groups M are, for example:

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The groups necessary for the novel compounds are linked to one another via bridges Y, such as O, S, COO, OCO, OCOO, CONH, NHCO, CON(R), N(R)CO or a direct bond, where R is C_1 - to C_4 -alkyl or hydrogen. Y is preferably O, OCO, COO or OCOO.

40

n is preferably 2.

Polymerizable groups which may be mentioned in particular are vinyl radicals, which are present, for example, in acrylic compounds, vinyl ethers and styrene derivatives. Epoxides are also suitable.

Examples of preferred radicals Z are the following:

,
$$=$$
 , $CH_2=CH-CH_2$, CO_2R

Z is particularly preferably CH₂=CH-, CH₂=C- or CP₂=CH-CH₂.

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- 15 The units $Z-Y[A]_m-Y-M-Y$ according to the invention, where Z, Y, m, A and M are as defined above, can be obtained by generally known synthetic methods, as described, for example, in EP-A 739403 and DE-A 39 17 196.
- 20 The chiral moieties can be purchased commercially and are thus available.

The novel compounds are suitable as high-twisting-power dopants for producing cholesteric liquid crystals. Using very small amounts, the reflection wavelength of the suitable state.

25 amounts, the reflection wavelength of the cholesteric mixture can be shifted into the visible region. It is advantageous here that the dopants form a stable mixture with the nematic host system, from which mixture the dopant does not crystallize out. The novel compounds are particularly suitable for this purpose.

The novel compounds are particularly suitable for use in electro-optical display elements or as chiral dopants for nematic or cholesteric liquid-crystal mixtures for producing layers which reflect in color.

The novel compounds are also suitable for use as chiral dopants in optical polarizers and optical filters.

Particular preference is given here to mixtures of the novel 40 compounds with nematic compounds of the formula

$$(Z)_{m}-Y-A-Y-M^{1}-Y-A-Y-(Z)_{m}$$
 II

where, in each case independently of one another, $45\ A,\ m,\ Y\ and\ Z\ are\ as\ defined\ above,\ and\ M^1$ is a mesogenic group.

Compounds of the formula II are described, for example, in DE-OS 195 32 408, GB-A-22 79 659, (EP-A-749 466) and EP-A-648 827.

5 Also claimed are solid compositions comprising the novel compounds in copolymerized form.

Also claimed are solid compositions comprising the novel compounds and nematic compounds of the above formula II, in each 10 case in copolymerized form.

Solid compositions of this type may be, for example, polymers which can contain partly uncrosslinked or non-copolymerized constituents. Non-copolymerized constituents of this type can be,

- 15 for example, unreactive organic molecules having a wide variety of molecular weights or alternatively inorganic or organic pigments. They may furthermore also be additives, for example rheology auxiliaries or light and/or oxidation stabilizers.
- 20 It is of course also possible for solid compositions of this type to be the polymerization products of the claimed liquid-crystalline mixtures containing the novel compounds, and of the claimed liquid-crystalline mixtures containing nematic compounds of the above formula II in addition to the novel compounds.

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Procedure:

3.84g (0.01 mol) of bis(4'-hydroxybenzoyl)-1,4:3,6-dianhydrosorbitol are dissolved in 30 ml of DMF and at 40°C for 3 hours and then at RT overnight. The reaction mixture is poured into water and acidified oxybenzoyl chloride are then added dropwise at 0 - 5° C. The mixture is stirred at $0-5^{\circ}$ C for one hour, 5.08 g (0.04 mol) of N,N-dimethylcyclohexylamine, and 8.0 g (0.024 mol) of 4-acryloxybutoxycarbonylusing conc. hydrochloric acid. Ethyl acetate is added, and the organic phase is washed with water, dried over sodium sulfate and evaporated in a rotary evaporator.

Yield: 11.3 g

The crude product is chromatographed over 1.1 kg of silica gel using the eluent petroleum ether/ethyl acetate (2:1).

Yield: 9.3 g (96%) [NMR]
HTP in ZLI 1840 63 µm⁻¹
Melting point: 109°C (DTA)
The compound has a right-handed twist.

Further compounds prepared by this and similar processes are the following:

HTP = helical twisting power

$$\sum_{R_1} - (CH_2)_{m^1} - Y_1$$

$$= \sum_{R_1} 0$$

			Т			Т	7		1							Г	Т	T	\neg
Twist	right-handed	The Part of the Pa	rignt-nanged	right-handed															
HTP mm ⁻¹ in ZLI 1840		50					·						63						
m ²	4	4	ۏ	9	4	2	2	4	9	4	2	2	4	9	9	4	•	4	9
m1	2	4	9	4	9	2	4	2	9	9	2	4	4	4	9	T T	-	4	9
R ²	Оме	ОМе	ОМе	ОМе	ОМе	н	H	H	H	H	H	H	æ	H	E	2	E	H	н
\mathbb{R}^1	ОМе	ОМе	ОМе	ОМе	Оме	æ	æ	H	H	H	H	1	I		=		Н	H	æ
¥2	0000	0000	0000	0000	0000	0000	0000	0000	0000	0000	0	C) C) C)		0	0000	0
Y 1	0000				0000		0000	0000	0000	0000	C						0000	0	0000
Ex.	2				9	7	8	6	6			: :	27	, ,	2 :	10	17	18	19

Twist		right-handed	right-handed	right-handed	right-handed	right-handed
HTP µm-1 in zi.t	1840					
m ²	y	0 4	, 4	9 4	. 9	
Im.	9	4	9	4	9	
R2	H	H	æ	В	H	
R1	×	æ	H	Н	Ξ.	
Y2	0000	000	000	0	2	
Y1	0	000		000		
ex .	21	22	23	24		

$$\begin{pmatrix} C(H_2)_m^1 - \gamma^1 \\ R^1 \end{pmatrix}$$

Β×.	Yl	Υ2	R1	R2	m	m ²	нтр	ms; at
					•		Lm-1))
							in ZLI 1840	
25	0000	0000	ОМе	ОМе	4	4	11	left-handed
26	0000	0000	ОМе	ОМе	9	9		left-handed
27	0000	0000	ОМе	ОМе	2	2		left-handed
28	0	0	Н	#	2	2		left-handed
29	0	0	н	н	4	2		left-handed
30	0	0	н	H	4	4	63	left-handed
31	0	0	Ħ	H	9	9		left-handed
32	000	000	н	н	2	2		left-handed
33	000	000	Н	H	7	4		left-handed
34	000	000	H	н	9	9		left-handed
35	000	000	H	н	9	4		left-handed
36	00 00	0000	H	н	2	2		left-handed
37	0000	೦೦೦೦	н	н	2	4		left-handed
38	0 00 0	0000	H	æ	4	4	13	left-handed
39	0000	0000	H	H	9	9		left-handed
40	0000	0000	н	н	4	9		left-handed
41	0000	0000	н	н	9	4		left-handed
42	0	0	ОМе	ОМе	4	4		left-handed

	R2	
-0-X1-X-0-	R1	

	R1				ONO	OGG	⊕ Σ	ÐW			17		
w2	- T		2		2		0		0000	000	220		
.		0		C	•	0			0000	000		000	
m ×		43		44		45	2.2	46	2	47		48	

Mixtures with liquid-crystalline substances

Example 49

5 x mol% of Example 1 are dissolved in the nematic mixture

by mixing the melts. After cooling to RT, cholesteric phases with the following reflection wavelengths - depending on X - are 25 obtained:

	x (mol%)	λ1 (nm)	_
	2.0	701	
30	3.0	482	
	4.0	341	

Example 50

x mol% of Example 1 are dissolved in the nematic compound 35

by mixing the melts. After cooling to RT, cholesteric phases with the following reflection wavelengths are obtained:

	x (mol%)	λ⊥ (nm)	
	3.05	660	
	3.51	565	
5	4.28	465	
	5.46 .	365	

Example 51

x mol% of Example 1 are dissolved in the nematic mixture

by mixing the melts. After cooling to RT, cholesteric phases with the following reflection wavelengths - depending on \mathbf{X} - are obtained:

30	x (mol%)	λ⊥ (nm)	
	2.0	854	
	2.5	696	
35	3.0 3.5	575	
	3.5	498	

Example 52

x mol% of Example 1 are dissolved in the nematic mixture

by mixing the melts. After cooling to RT, cholesteric phases with the following reflection wavelengths - depending on X - are 25 obtained:

	x (mol%)	λ⊥ (nm)	_
	1.5	974	
	2.0	736	
30	2.5	588	
	3.0	493	

Comparative Example 53

 $_{4 \text{ mol}\$}$ of both the compound from Example 1 and the compound of the formula III (corresponding to EP-A-739 403)

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are added to the nematic compound

20 The two substances are melted and their spectrum measured at RT. The compound from claim 1 gives a reflection at 431 nm, while the compound of Formula III gives a reflection at 1087 nm.

<u>Claims</u>

1. A compound of the formula

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15

$$(Z-Y-[A]_m-Y-M-Y-)_n X$$

I,

where, independently of one another,

10 A is a spacer,

which are unsubstituted or substituted by C₁- to C₄-alkyl, methoxy, ethoxy, fluorine, chlorine, bromine, C₁- to C₂₀-alkoxycarbonyl or C₁- to C₂₀-alkylcarbonyl and are linked via 0, CO, OCO, OCOO or COO,

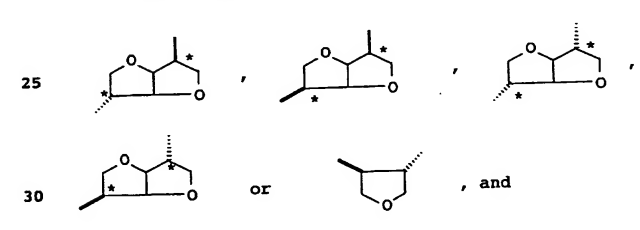
Y is a direct bond, O, S, COO, OCO, OCOO, CON(R) or N(R)CO, and

z is a polymerizable group,

m is 0 or 1,

20 n is from 2 to 6, and

X is a chiral radical of the formula



35 R is C_1 - to C_4 -alkyl or hydrogen.

- 2. A compound as claimed in claim 1, where n = 2.
- 3. A compound as claimed in claim 1, where M is

- 10 4. A compound as claimed in claim 1, where
 Y is 0, 0CO, COO, 0COO or a direct bond.
- 5. A compound as claimed in claim 1, where

Z is
$$CH_2=CH$$
—, CH_2 — CH_3 or $CH_2=CH$ — CH_2 .

20

6. A compound as claimed in claim 1, where

25 A is
$$(CH_2)_p$$
, $(CH_2CH_2O)_qCH_2-CH_2$, $(CH_2CH_2O)_q-CHCH_2$ CH3 CH3 CH3 CH3

- p is from 1 to 12 and q is from 1 to 3.
 - 7. A compound as claimed in claim 1, where

- A compound as claimed in claim 1 and referred to in any 8. of the foregoing Examples 1 to 52.
- A liquid-crystalline mixture comprising one or more compounds as claimed in any of claims 1 to 8. 9.
- A liquid-crystalline mixture comprising one or more compounds as claimed in any of claims 1 to 8 and one or 5 10. more nematic compounds of the formula II

II $(Z)_{m}-Y-A-Y-M^{1}-Y-A-Y-(Z)_{m}$

- where, in each case independently of one another, A, m, Y and Z are as defined in claim 1, and M^1 is a mesogenic 10 group.
 - A solid composition comprising a compound as claimed in 11. any of claims 1 to 8 in copolymerized form.
- A solid composition comprising a compound as claimed in any of claims 1 to 8 and a nematic compound of the 12. formula II defined in claim 10, in each case in 15 copolymerized form.
- The use of a compound as claimed in any of claims 1 to 8 in electro-optical displays or as a chiral dopant for 13. nematic or cholesteric liquid crystals for producing 20 layers which reflect in color.
 - The use of a compound as claimed in any of claims 1 to 8 as a chiral dopant in optical polarizers and optical 14. filters.





Application No:

GB 9821817.5

Claims searched: 1-14

Examiner:

Stephen Quick

Date of search: 8 January 1999

Patents Act 1977
Search Report under Section 17

Databases searched:

UK Patent Office collections, including GB, EP, WO & US patent specifications, in:

UK Cl (Ed.Q): (not searched)

Int Cl (Ed.6): (not searched)

Other: Online: CAS ONLINE

Documents considered to be relevant:

Category	Identity of documen	nt and relevant passage	Relevant to claims
A	WO 95/16007 A1	(BASF), see pages 3 (lines 1-11), 4 (lines 40-end), 5 (lines 1-17) & 8 (lines 33-42); acknowledged in this application as EP 0739403	-

& Member of the same patent family

- Document indicating technological background and/or state of the art.
- P Document published on or after the declared priority date but before the filing date of this invention.
- E Patent document published on or after, but with priority date earlier than, the filing date of this application.

X Document indicating lack of novelty or inventive step

Y Document indicating lack of inventive step if combined with one or more other documents of same category.